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Dynamic changes of gut microbiota and hepatic functions are different among biliary atresia patients after Kasai portoenterostomy

Dear Editor,

Biliary atresia (BA) is a fatal neonatal disease resulting in cholestasis and progressive hepatic failure. Despite the fact that Kasai portoenterostomy (KP) can restore bile drainage, impaired hepatic functions and even advanced cirrhosis can be detected in most of the patients. Given the interaction between gut microbiota (GM) and hepatic functions, this study aimed to assess if BA patients with different pre-surgery GM had different GM dynamics and hepatic functions following KP.

According to the inclusion criteria (see the Supporting Information), we enrolled 26 BA patients aged 19–105 days whose parents approved KP therapy. Then we collected feces for sequencing of 16S rDNA V3-V4 regions at five timepoints (Figure S1A). Before KP, 18 patients had Bifidobacterium-dominated (BD) and eight patients had non-Bifidobacterium-dominated (NBD) GM structure (Figure 1A,B). In the NBD group, Bacteroides, Rothia, Defluviiitoga and Collinsella represented higher relative abundance when compared to the BD group (Figure 1A). Additionally, microbial samples in the BD group had significantly lower diversity than the NBD group (Figure 1C). Further analysis indicated that inter-individual dissimilarity in the BD group was dramatically lower than that in the NBD group (Figure 1D). In addition, the inter-individual distance between BD and NBD samples was as high as that in the NBD group (Figure 1D). At the phylum level, Actinobacteria represented obviously higher relative abundance in microbial samples of the BD group (averaged 92.99% vs. 49.92% in the NBD group) (Figure 1E). Nonetheless, the average level of Bacteroidetes and Firmicutes in the NBD group was 3.54 and 10.98 times as that in the BD group, respectively (Figure 1E). Functional prediction indicated accumulation of amino acid metabolism, depletion of cell motility, cellular processes and signalling in microbial samples of the BD group (Figure S1B). Intravenous antibiotic exposure caused slight but insignificant changes to the GM composition of BD and NBD patients (Figure S1C).

Permutational multivariate analysis of variance (PERMANOVA) unravelled that KP contributed remarkably to post-surgery GM changes (Figure 2A). Consistently, GM changed significantly after KP (Figure S1D). Inter-group distance of microbial samples kept high even after KP, which suggested distinct GM dynamics between BD and NBD groups (Figure 2B,C,D). Consistent with transition of Bifidobacterium-dominated GM structure to other GM types following development, microbial samples in the BD group had higher dynamic changes than those in the NBD group after KP (Figure 2B). A previous study also found a decreased level of Bifidobacterium in patients who had clearance of jaundice. Inter-individual differences of post-surgery GM changes in the BD group may be explained by different sensitivities of Bifidobacterium strains to bile.

Of the microbial samples in the BD group, phylum Actinobacteria levels reduced, while Bacteroidetes and Firmicutes levels elevated after KP (Figure 2E). There was an opposite trend for dynamic changes of Actinobacteria, Bacteroidetes and Firmicutes in the NBD group (Figure 2E). After surgery, contributions of the bacterial genus in inter-group differences also changed (Figure 2F). For instance, Bifidobacterium was the most robust in contributing to differences between BD (mean ± SD, 88.32 ± 12.87%) and NBD (29.46 ± 10.61%) groups before KP, and was insignificant in explaining inter-group differences (BD: 37.65 ± 32.31%, NBD: 40.18 ± 31.35%) after therapy (Figure 2F). By contrast, the significance of Bacteroides in contributing to inter-group differences increased after KP (BD: 17.07 ± 29.43%, NBD: 9.24 ± 24.00%) (Figure 2F). This possibly suggested normal development of Bifidobacterium-dominated GM to Bacteroides-dominated enterotype. Additionally,
Defluviitoga and Fusobacterium kept significant contributions to inter-group differences both before and after KP (Figure 2F). In the early stage of life, Bifidobacterium dominated GM, and depletion of Bifidobacterium correlated with a variety of diseases. This may explain better hepatic functions for patients in the BD group before surgery (Figure 3A). PERMANOVA indicated different impacting factors for clinical indicators (Figure 3B). Like GM (Figure 2B), dissimilarity of clinical indicators at different time points was slightly higher in the BD group than in the NBD group (Figure 3C). Despite similar changes following KP, ALT, AST, IBIL levels kept higher in the NBD group and TAB levels kept higher in the BD group (Figure 3D), suggesting the long-term impact of pre-surgery GM on hepatic functions. Correlation analysis indicated the association of amino acid metabolism, nucleotide metabolism, membrane transport and signalling molecules and interaction with Bifidobacterium, which decreased after KP (Figures 2F and 3E). Nevertheless, environmental adaptation, signal transduction, lipid and energy metabolism correlated with bacterial genus, which had high importance in inter-group differences after KP, including Defluviitoga, Fusobacterium and Bacteroides (Figures 2F and 3E).

Further analysis found negative correlation of Akkermansia, Parabacteroides, Actinomyces, Fusobacterium and Finegoldia with serum levels of bilirubin, while Prevotella correlated negatively with AST and ALT levels in the BD group (Figure S1E). This was partly consistent with prior reports suggesting that Prevotella was associated with short fatty acids metabolism and Akkermansia was a promising probiotics candidate.9 In the NBD group, serum levels of bilirubin, AST and ALT correlated negatively with Bifidobacterium and Collinsella, but positively correlated with...
FIGURE 2  Post-surgery gut microbiota (GM) dynamics after surgery. (A) PERMANOVA to assess the effects of several indices on GM dynamics: more significant if the point is located near the outer circle. Kasai portoenterostomy (KP) (p-value = .002), time (p-value = 0.044) and pre-surgery GM structure (“Group” in the figure, p-value = 0.048) contributed remarkably to post-surgery GM changes. Permutation test was applied to assess the statistical significance. (B) Bray-curtis distance (dissimilarity) of microbial samples between different timepoints for *Bifidobacterium*-dominated (BD) and non-*Bifidobacterium*-dominated (NBD) groups. (C) Comparison of dynamic Bray-curtis distance (dissimilarity) between microbial samples in BD and NBD groups, respectively, as well as between BD and NBD groups. (D) Alpha diversity of microbial samples at different timepoints for BD and NBD groups. (E) Phylum-level microbial components at different timepoints for BD and NBD groups. (F) Inter-group genus differences: the left sub-figure represented differences before surgery (microbial samples collected before KP), the right sub-figure represented differences after surgery (all microbial samples collected 1, 3, 7, 30 days after KP), and the importance decreased from top to bottom (e.g., *Bifidobacterium* represents the most importance in inter-group differences before KP). BD: *Bifidobacterium*-dominated GM structure before KP; NBD: non-*Bifidobacterium*-dominated GM structure before KP. Time 1, 2, 3, 4 and 5 represent the day of hospitalization (before surgery and intravenous antibiotics exposure) 1, 3, 7 and 30 days after surgery, respectively. Error bars represent the standard error of the mean. Wilcoxon rank-sum test was applied to analyse statistical significance for inter-group differences. *p-value < .05, **p-value < .01, ***p-value < .001, ****p-value < .0001
FIGURE 3  Different dynamic changes of several clinical indicators between *Bifidobacterium*-dominated (BD) and non-*Bifidobacterium*-dominated (NBD) groups. (A) Difference of clinical indicators between BD and NBD groups before surgery. AST, IBIL, and TBIL were statistically increased in NBD patients compared to BD (304.89 ± 237.91 vs. 221.36 ± 115.46 U/L; 59.97 ± 41.48 vs. 42.27 ± 17.69 μmol/L; 190.99 ± 66 vs. 182.56 ± 60 μmol/L in the BD group, respectively), and ALT levels were increased but not statistically significant.
opportunistic pathogens *Saccharopolyspora*, *Corynebacterium* and *Defluviitoga* (Figure S1E). Additional analysis indicated the involvement of clinical indicator-related microbial genus in the co-occurrence network (Figure S1F). For example, *Akkermansia* and *Fusobacterium* were in positive correlation in the BD group, while *Saccharopolyspora* and *Defluviitoga* were in positive correlation in the NBD group (Figure S1F). The above-mentioned correlations implicated the necessity to consider pre-surgery GM structure when applying prebiotics or probiotics to improve hepatic functions after KP.10

Though this study found different dynamic changes of GM and clinical indicators for BA patients who received KP therapy, several limitations should be noted. First, 16S rRNA gene amplicon sequencing only analysed known bacterial genus. Additionally, GM-derived metabolites were not detected to analyse the interaction between GM and hepatic functions. Second, we only performed sample collection in 30 days after surgery, thus it was impossible to assess long-term effects of pre-surgery GM on post-surgery hepatic functions. Finally, we did not record factors that affected early-life GM, including delivery mode and feeding pattern. However, lack of information like delivery mode will not affect our findings negatively because the aim was to assess the association of developed GM with hepatic functions.

In conclusion, our study identified different hepatic functions for BA patients with different pre-surgery GM structures, and emphasised the importance of pre-surgery GM structures in GM dynamics and restoration of hepatic functions after receiving KP therapy.

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**CONFLICT OF INTEREST**

The authors declare that they have no conflict of interest.

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